#### (19) World Intellectual Property Organization International Bureau





(43) International Publication Date 18 May 2006 (18.05.2006)

# (10) International Publication Number WO 2006/051387 A1

(51) International Patent Classification: C12N 1/16 (2006.01) C12N 9/02 (2006.01) C07K 14/37 (2006.01) C12Q 1/34 (2006.01)

(21) International Application Number:

PCT/IB2005/003352

(22) International Filing Date:

9 November 2005 (09.11.2005)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data: 2004/9060

9 November 2004 (09.11.2004) ZA

- (71) Applicant (for all designated States except US): UNIVER-SITY OF STELLENBOSCH [ZA/ZA]; Victoria Street, 7600 Stellenbosch (ZA).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): BAUER, Florian, Franz [ZA/ZA]; 3 Serruria Street, Paradyskloof, 7600 Stellenbosch (ZA). SWIEGERS, Jan, Hendrik [ZA/ZA]; 2755 Lipkin Street, 7141 Betty's Bay (ZA).
- (74) Agents: GILSON, David, Grant et al.; P.O. Box 454, 0001 Pretoria (ZA).

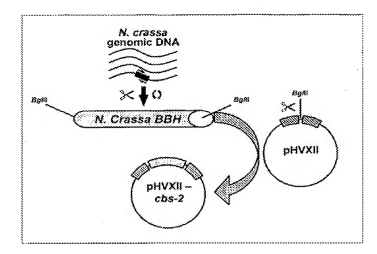
- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
- (84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, FT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

#### Published:

- with international search report
- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments

[Continued on next page]

(54) Title: METHOD OF PRODUCING A CARNITINE-SYNTHESISING MICRO-ORGANISM



(57) Abstract: The invention discloses a method of producing a micro-organism that can biosynthesise carnitine from a non-carnitine synthesising micro-organism, the method including the step of transforming the non-carnitine synthesising micro-organism with a nucleotide sequence encoding  $\gamma$ -butyrobetaine hydroxylase (BBH). The transformed micro-organism is capable of producing carnitine when cultured in the presence of gamma-butyrobetaine. A method of identifying a carnitine-producing micro-organism is also disclosed, the method including the steps of applying a micro-organism to a synthetic agar medium which does not contain carnitine and is coated with a layer of Saccharomyces cerevisiae  $\Delta cit2$  strain; culturing the micro-organism; and detecting the presence of a zone in the agar medium formed by carnitine-producing micro-organisms. The transformed micro-organism may be used to enhance the nutritional value of foods and beverages, such as bread and beer.



## WO 2006/051387 A1

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

#### METHOD OF PRODUCING A CARNITINE-SYNTHESISING MICRO-ORGANISM

5

15

20

25

## BACKGROUND OF THE INVENTION

The invention relates to a method for producing a micro-organism that is capable of synthesising carnitine, and also describes a method of identifying whether a microorganism is capable of synthesising carnitine.

L-Carnitine (3-hydroxy-4-*N*-trimethylaminobutyrate) is a quaternary ammonium compound that was first discovered in muscle extracts in 1905 (Bremer, 1983). In 1952, it was shown that the mealworm *Tenebrio molitor* is dependent on carnitine for survival, generating new interest in this molecule, which was named vitamin  $B_T$  (Carter et al., 1952). Later investigations showed that most eukaryotic organisms could synthesise L-carnitine from trimethyllysine as a precursor (Vaz and Wanders, 2002). Nevertheless, carnitine deficiencies occur and they are debilitating diseases, frequently due to genetic mutations (Bonnefont et al., 1999; Lahjouji et al., 2001). Such diseases are characterised by low levels of carnitine in either the serum or in specific tissues. In most cases, patients respond favourably to exogenous dietary supplementation of carnitine (Pons and De Vivo, 1995). In recent times, L-carnitine has also been used for symptomatic treatment in cases of diseases, such as chronic fatigue syndrome, coronary vascular disease, hypoglycemia and muscular myopathies (Kelly, 1998). In addition, carnitine is widely used in nutritional products, such as energy drinks, weight loss supplements and baby formulae (Carter et al., 1995).

In mammalian cells, carnitine is an essential component of the mitochondrial carnitine cycle that is responsible for the transfer of activated long-chain fatty acids into the mitochondria or peroxisome for β-oxidation (Bieber, 1988). In the yeast Saccharomyces cerevisiae, on the other hand, β-oxidation occurs solely in the peroxisomes (Kunau et al., 1988). Van Roermund et al. (1995) showed that exogenous carnitine was essential for growth on fatty acids as sole carbon source in

the absence of the glyoxylate cycle citrate synthase, Cit2p. Later, Swiegers et al. (2001) showed that in the  $\Delta cit2$  strain, carnitine is essential for growth on all non-fermentable carbon sources. Therefore, S. cerevisiae is unable to biosynthesise carnitine endogenously but relies on exogenous carnitine, which is transported into the cell by the general amino acid membrane transporter Agp2p (van Roermund et al., 1999; Swiegers et al., 2001).

Mammals, plants and some fungi are able to biosynthesise carnitine from E-Ntrimethyllysine (TML) (Lindstedt and Lindstedt, 1970; Kaufman and Broquist, 1977; Bremer, 1983). In mammals, TML is provided by the lysosomal hydrolysis of proteins that contain this amino acid as a result of the post-translational modification of lysine residues (Bremer, 1983). However, in Neurospora crassa, free lysine is trimethylated in the cytosol (Borum and Broquist, 1977). In the first step of carnitine biosynthesis, TML is hydroxylated to β-hydroxy-ε-N-trimethyllysine by ε-N-trimethyllysine hydroxylase (TMLH; EC1.14.11.8) (Rebouche and Engel, 1980; Bremer, 1983). Subsequently, βhydroxy-e-N-trimethyllysine is cleaved into y-trimethylamino-butyraldehyde and glycine by β-hydroxy-ε-N-trimethyllysine aldolase (Rebouche and Engel, 1980; Bremer, 1983). The aldehyde is then oxidised by y-trimethylaminobutyraldehyde dehydrogenase to form γ-butyrobetaine (Hulse and Henderson, 1980; Rebouche and Engel, 1980; Finally, y-butyrobetaine is hydroxylated at the 3-position by Bremer, 1983). γ-butyrobetaine hydroxylase to form L-carnitine (Figure 1) (BBH; EC 1.14.11.1) (Englard, 1979; Rebouche and Engel, 1980; Bremer, 1983).

The identity of some of the intermediate metabolites of the carnitine biosynthesis pathway was first elucidated in the filamentous fungus *Neurospora crassa*, using isotope-labelling experiments (Kaufman and Broquist, 1977). The genes encoding the enzymes required for the catalysis of three of the four reactions required for carnitine biosynthesis have been characterised at the molecular level, in rats and humans (Vaz et al., 1998; Galland et al., 1999, Vaz et al., 2000; Vaz et al., 2001).

The applicant has therefore identified a need for producing a strain of *S. cerevisiae* that can produce carnitine. The applicant has also identified a need for a simple and inexpensive method of determining whether or not a micro-organism is capable of producing carnitine.

5

10

15

20

25

30

#### **SUMMARY OF THE INVENTION**

According to a first embodiment of the invention, there is provided a method of producing a micro-organism that can biosynthesise carnitine from a non-carnitine synthesising micro-organism, the method including the step of:

transforming the non-carnitine synthesising micro-organism with a nucleotide sequence encoding y-butyrobetaine hydroxylase (BBH).

10 The micro-organism may be a yeast strain, such as Saccharomyces cerevisiae.

The transformed strain may be cultured in the presence of  $\gamma$ -butyrobetaine in order for the strain to produce carnitine.

The γ-butyrobetaine hydroxylase may be a *Neurospora crassa* γ-butyrobetaine hydroxylase, and the nucleotide sequence may be a genomic fragment having the nucleotide sequence as set out in SEQ ID NO: 1 (Figure 7).

The S. cerevisiae strain may be a FY23 strain.

20

25

35

5

The non-carnitine synthesising micro-organism may be identified by:

applying the micro-organism to a synthetic culture medium containing a non-fermentable carbon source and no carnitine, and which is coated with a layer of Saccharomyces cerevisiae \( \Delta cit2 \) strain;

culturing the micro-organism; and

detecting formation of a zone produced in the culture medium in the region of the micro-organism if it is a carnitine-producing micro-organism, or detecting no such zone if the micro-organism is not able to produce carnitine.

30 The above identification steps may also be performed to determine whether the method of producing a micro-organism that can biosynthesise carnitine has been successful.

According to a second embodiment of the invention there is provided a strain of Saccharomyces cerevisiae that can produce carnitine in the presence of  $\gamma$ -

butyrobetaine, which has been transformed with a nucleotide sequence encoding  $\gamma$ -butyrobetaine hydroxylase (BBH) substantially as described above.

According to a third embodiment of the invention, there is provided a method of identifying a carnitine-producing micro-organism, the method including the steps of:

applying a micro-organism to a synthetic culture medium containing a non-fermentable carbon source, no carnitine and coated with a layer of Saccharomyces cerevisiae \( \Delta cit 2 \) strain;

culturing the micro-organism; and

detecting formation of a zone produced in the culture medium in the region of the micro-organism if the micro-organism is a carnitine-producing micro-organism.

The culture medium may be illuminated or transilluminated in order to detect formation of the zone.

The culture medium may be agar, and the non-fermentable carbon source may be ethanol. For example, the culture medium may be 2% ethanol synthetic agar medium.

The micro-organism may be cultured for a period of about 10 days at about 30 °C.

20

5

10

15

According to a fourth embodiment of the invention, there is provided a method of producing carnitine, the method including the step of culturing in the presence of y-butyrobetaine hydroxylase (BBH) a micro-organism that has been transformed as above.

25

30

The micro-organism may be included in the production of a beverage or food product, such as beer or bread.

According to a fifth embodiment of the invention, there is provided a method of enhancing the nutritional value of a beverage or food product, the method including the step of incorporating a microorganism that has been transformed as described above in the process of manufacturing the food or beverage product.

#### **BRIEF DESCRIPTION OF THE DRAWINGS**

Figure 1 shows a reaction scheme for hydroxylation of γ-butyrobetaine to L-carnitine by γ-butyrobetaine hydroxylase;

5

15

20

25

Figure 2 shows the alignment of the 671 amino acids expressed *N. crassa* BBH homologue (SEQ ID NO: 2) to human, *Caenorhabditis elegans* and *Pseudomanos* BBH proteins;

10 Figure 3

shows a carnitine large-scale plate screen. (A) Strains were grown on glucose synthetic medium and streaked on a 2% ethanol synthetic medium agar plate with 10 mg/l y-butyrobetaine and with a thin mat of Δcit2 cells grown on synthetic glucose medium, which were washed twice with sterile distilled water before plating. Cells were grown for 10 days at 30°C. The production of L-carnítine by the strain expressing a functional BBH results in the secretion of carnitine, which complements the surrounding  $\Delta cit 2$  strains and resulted in the formation of a zone. (B) Carnitine secretion plate assay for identification of endogenous biosynthesis and secretion of carnitine. Yeasts were grown on glucose synthetic medium before they were streaked onto a 2% ethanol synthetic medium agar plate with a thin mat Δcit2 cells. Cells were grown for 10 days at 30°C. Yeast strains with endogenous L-carnitine biosynthesis and secretion could be identified by the zone formation (D5, D4 and D2). (D5) Yarrowia lipolytica; (D4) Rhodotorula graminis; (D2) Candida curvata. The controls were the S. cerevisiae laboratory strain FY23 (A1) and industrial commercial wine strain VIN13 (A2). Other non-zone forming yeast represents a various collection of different genus and To improve visual detection of zones, plates were species. transilluminated with light and photos taken. Biosynthesis of carnitine in zone forming strains was verified using electrospray mass spectrometry as described below;

30

35

Figure 4 shows (A) photographs of petri dishes on which strains were grown for 4 days at 30°C on synthetic glycerol (3%) medium and synthetic glycerol medium with 10 mg/l y-butyrobetaine; and (B) growth curves of strains

and transformants: FY23 wild type strain ( $\blacktriangle$ ); FY23 $\triangle$ cit2 strain ( $\triangle$ ); FY23 wild type strain with cbs-2 ( $\blacksquare$ ); FY23 $\triangle$ cit2 strain with cbs-2 ( $\square$ ). Each strain was grown in 100 ml of synthetic glycerol (3%) medium plus 10 mg/l  $\gamma$ -butyrobetaine at 30°C;

5

Figure 5 shows the measurement of intracellular carnitine and acetylcarnitine

using electrospray mass spectrometry. (A) FY23 wild type strain with cbs-2 and (B) FY23 wild type strain was grown on synthetic glycerol (3%) medium with 10 mg/l  $\gamma$ -butyrobetaine for 4 days at 30°C, after which cells were harvested and intracellular carnitine and acetylcarnitine levels were determined. Carnitine has a parent ion of 162 and the daughter fragment of 43 was measured. Acetylcarnitine has a parent ion of 204 and a daughter ion of 85 was measured. The level of intracellular carnitine measured for cbs-2 transformed cells were 897 ng/gWW and acetylcarnitine 1151 ng/gWW;

15

10

Figure 6 shows a diagrammatic representation of the cloning strategy employed in the construction of the *N. crassa* BBH containing construct pHVXII-cbs-2. The symbol "≫" indicates the use of restriction enzymes, and "♣" refer to the use of the polymerase chain reaction;

20

Figure 7 shows the nucleotide sequence of a 2016bp fragment (SEQ ID NO: 1), encoding a putative protein with BBH homoloy, cloned from *N. crassa* genomic DNA in FASTA format; and

25

Figure 8 shows the putative amino acid sequence (SEQ ID NO: 2) encoded by the pHVXII-cbs-2 construct, which includes the entire area of BBH homology, in FASTA format.

30

## DETAILED DESCRIPTION OF THE INVENTION

A method of producing a micro-organism that can biosynthesise carnitine from a noncarnitine synthesising micro-organism is described herein. The method includes the

step of transforming the non-carnitine synthesising micro-organism with a nucleotide sequence encoding  $\gamma$ -butyrobetaine hydroxylase (BBH).

The micro-organism is generally a yeast strain, such as  $Saccharomyces\ cerevisiae$ , and more particularly a FY23 strain, although it may be any other type of micro-organism that is able to transport butyrobetaine into the cell, as the transformed strain is cultured in the presence of  $\gamma$ -butyrobetaine in order for the strain to produce carnitine.

The γ-butyrobetaine hydroxylase is generally a *Neurospora crassa* γ-butyrobetaine hydroxylase, and the nucleotide sequence thereof may be a genomic fragment having the nucleotide sequence as set out in SEQ ID NO: 1 (Figure 7).

Also disclosed is a method of identifying a carnitine-producing micro-organism, by applying a micro-organism to a synthetic culture medium which contains a non-fermentable carbon source but no carnitine, and which is coated with a layer of *Saccharomyces cerevisiae* Δ*cit2* strain. A typical culture medium is agar, and a suitable non-fermentable carbon source is ethanol. For example, the culture medium may be 2% ethanol synthetic agar medium. The micro-organism is then cultured, typically for a period of about 10 days at about 30 °C. The formation of a zone produced in the culture medium in the region of the micro-organism will be detectable, such as by illumination or transillumination, if the micro-organism is a carnitine-producing micro-organism. No such zone will be detected if the micro-organism is not a carnitine-producing micro-organism.

25

30

20

15

5

L-Carnitine is a quaternary ammonium compound that plays an essential role in the transfer of activated acyl-residues across intra-cellular membranes. Most eukaryotes can neo-synthesise carnitine, but recent data show that this is not the case in the yeast  $Saccharomyces\ cerevisiae$ . The filamentous fungus  $Neurospora\ crassa$  was one of the first organisms used to identify the precursor and intermediates of the carnitine biosynthesis pathway. In this organism, the precursor trimethyllysine is converted in a four-step process to carnitine. In the last step of this pathway,  $\gamma$ -butyrobetaine is hydroxylated to form carnitine in a reaction catalysed by  $\gamma$ -butyrobetaine hydroxylase (BBH).

A novel plate screen that can be used to identify genomic fragments of *N. crassa* that functionally express BBH is described herein. Using this plate screen, a genomic fragment encoding the *N. crassa*  $\gamma$ -butyrobetaine hydroxylase (BBH) was identified and the gene designated *cbs-2*. The invention teaches that a wild type yeast strain transformed with the *cbs-2* gene can use exogenous  $\gamma$ -butyrobetaine to produce carnitine, and expression of this gene is able to rescue the growth defect of a  $\Delta cit2$  strain on non-fermentable carbon sources, without carnitine, in the presence of  $\gamma$ -butyrobetaine.

10

5

The invention will now be described in more detail by way of the following non-limiting examples.

#### Examples:

15

20

25

30

#### Yeast strains and plasmids

FY23 (MATa leu2 trp1 ura3) was used as a wild-type strain (Winston et al., 1995). The FY23Acit2 (MATa leu2 ura3 cit2::TRP1) was used as the glyoxylate citrate synthase deficient strain (Swiegers et al., 2001). A 2016 bp fragment (Figure 7; SEQ ID NO: 1) was cloned from N. crassa genomic DNA using the primers NcBBH-F (5'-GATCAGATCT ATG AAA GTC GAC AAG GAA GCC GGC AA-3') (SEQ ID NO: 3) and NoBBH-R (5'-GATCAGATCT TTA TGC GTT CCA GTT CAC CGT GCC CAA-3') (SEQ ID NO: 4) with introduced restriction sites. Genomic DNA was extracted from strain PPRI 3338 (National Collection of Fungi, Agricultural Research Council, Pretoria, South Africa). The fragment was cloned into expression vector pHVXII into the Bg/II site under the regulation of the PGK1 promoter (Volschenck et al., 1997; represented in Figure 6). Sequencing was done using the ABI-Prism sequencer. The S. cerevisiae gene YHL021c was amplified by PCR from genomic DNA from strain FY23 using the primers YHL-F (5'-GATCGAATTC ATG CTA AGA TCA AAT TTA TGC AGA GGA-3') (SEQ ID NO: 5) and YHL-R (5'-GATCCTCGAG TTA TTT GTA CTG AGG AAA CTT CTC TTC-3') (SEQ ID NO: 6) with introduced restriction sites. The fragment was cloned into expression vector pHVXII into the BallI site under the yeast PGK1 promoter. Constructs were transformed into the yeast strains using the lithium acetate procedure (Becker and Gaurente, 1991).

## Media and growth conditions

5

10

15

20

35

Escherichia coli carrying plasmids were grown in Luria Bertani (LB) broth with 10 mg/l ampicillin. Yeast strains were grown in YPD (1% yeast extract, 2% bactopeptone, 2% glucose), synthetic glucose medium (6.7 g/l yeast nitrogen base without amino acids, 2% glucose, amino acids as required), and synthetic glycerol medium (6.7 g/l yeast nitrogen base without amino acids, 3% glycerol, amino acids as required). Media was prepared using double distilled water.

## Intra-cellular carnitine extraction

Transformants were grown on synthetic glucose medium for two days and then inoculated in 100 ml synthetic glycerol medium with 10 mg/l y-butyrobetaine and grown for 4 days at 30°C. Cells were harvested by centrifuging 5 min at 5000 rpm and washed with 40 ml double distilled water and harvested again using the same procedure. Cells were resuspended in 1 ml double distilled water, transferred to a 1.5 ml microcentrifuge tube and harvested at 12 000 rpm for 2 min. Wet weight was determined by weighing the cells and the microcentrifuge tube after all the supernatant was removed by pippeting. The cells were resuspended in 0.2 ml double distilled water. The cells were disrupted by adding 0.16 g glass beads and vortexed for 30 min at 8°C. The cells were then vortexed for 10 min at 12 000 rpm and 0.1 ml of the supernatant added to 0.9 ml acetonitrile and stored at -20°C. Before ES-MS analysis, the solution was centrifuged for 10 min at 12 000 rpm to remove all protein precipitates and the supernatant used for analysis.

## HPLC-electrospray mass spectrometry

Mass spectrometry was performed on a Micromass (Manchester, UK) Quattro triple 25 quadropole mass spectrometer fitted with an electrospray ionisation source. Solvent A (acetonitrile/water/formic acid: 30/70/0.05 (v/v/v) was used as a carrier solvent and was supplied to the ionisation source by a LKB/Pharmacia (Sweden) pump. For direct injection of the carnitine and acetylcarnitine standards, the flow rate was 20 µl/min and 5 μl of the standard was injected through a Rheodyne injection valve. The molecular 30 ion ([M+H]+) of carnitine and acetylcarnitine was observed using a capillary voltage of 3.5 kV, source temperature of 80°C and a cone voltage setting of 20 V. To obtain the fragment pattern of carnitine and acetylcarnitine, the molecular ion was dissociated in the fragmentation cell by collision-induced dissociation at an argon pressure of 2.8x10<sup>-3</sup> mbar (2.8x10<sup>2</sup> Pa) applying collision energy of 20 eV. The resultant fragments

were scanned in the second analyser. Quantitation of carnitine and acetylcarnitine in the incubation samples was accomplished by LCMSMS. A Luna C18 150x2 mm (3µ) column was used for separation, with solvent A as the mobile phase at a flow rate of 100 µl/min delivered by the above mentioned pump. Five microliter of sample was injected by a Waters 747 autosampler. The eluent from the column was directed into the electrospray ionisation source of the mass spectrometer. The capillary voltage, cone voltage, argon pressure and collision energy were as mentioned above. Detection was by multiple reaction monitoring, using the molecular ions of carnitine and acetylcarnitine as precursor ions and the fragments at m/z = 43 and m/z = 85 as product ions, respectively. Chromatographic peaks representing carnitine and acetylcarnitine were integrated and the concentration in the incubation samples were calculated from a dilution range of known concentrations of standard carnitine and acetylcarnitine in distilled water and diluted to a final concentration of 90/10 (v/v): acetonitrile/15 mM Tris.HCl. The calculations were automatically performed by the Quantify program of MassLynx and expressed as ng/ml.

## Identification of a N. crassa BBH homologue

5

10

15

20

35

BBH protein sequences from different organisms are highly homologous to each other and to TMLH protein sequences. BBH and TMLH are part of a family of  $\alpha$ ketoglutarate-dependent, non-haem ferrous iron dioxygenases (Vaz and Wanders, 2002). However, when the BBH and TMLH proteins are compared to other proteins using BLAST searches (NCBI), reduced homology is found, indicating that these enzymes form a separate class of dioxygenases.

Searching the N. crassa Genome Database (NCGD) resulted in the identification of 2 25 putative proteins with high homology to human, rat and mouse BBH protein sequences (http://www-genome.wi.mit.edu/annotation/fungi/neurospora/). The first corresponds to a TMLH previously cloned and identified in a laboratory at Stellenbosch University (Swiegers et al., 2002), whereas the second was a novel gene encoding a hypothetical protein (NCU06891.1). The BBH homologous gene predicted by the NCGD consists of 30 5 predicted exons totalling 3786 bp and translating a hypothetical protein of 1262 aa (http://www-genome.wi.mit.edu/annotation/fungi/neurospora, Feature Search: NCU06891.2; SEQ ID NOS: 8 and 9). This is in strong contrast to the other known BBH proteins of humans, rats, mice and Pseudomonas, the length of which varies between 340-380 aa. However, homology to BBH proteins is only found for the protein

sequence translated by the last exon (exon 5) as described in the feature map of the hypothetical protein on the NCGD. The other translated exons do not show homology to any known protein.

Using the *N. crassa* genomic DNA, a 2016 bp fragment (Figure 7, SEQ ID NO: 1) (encoding a putative protein of 671 aa which includes the entire area of BBH homology, Figure 8 (SEQ ID NO: 2)), was cloned into a yeast expression vector, pHVXII under regulation of the *PGK1* promotor. Sequencing confirmed that the correct genomic area was cloned. Homology of the 671 aa putative protein to other known BBH proteins from humans, mouse and *Pseudomonas* are shown in Figure 2. However, the 671 aa putative proteins contained a 111 aa N-terminal and 110 aa C-terminal flanking regions without any homology to known BBH proteins. The C-terminal domain contains a sixfold repeat of the sequence "PKVEE" (SEQ ID NO: 7). Some 'additional' internal sequences, which contained GGGG repeats, were also present within the BBH homologous area, similar to what was observed for the *N. crassa* TMLH where an 11 residue poly P region and an "AAAAA" are found within the TMLH homologous area (Swiegers et al., 2002).

#### Screening of carnitine producing transformants

5

10

15

20

25

30

35

A large-scale screen was developed to identify microorganisms producing carnitine. The screen is based on the carnitine-dependent Δ*cit2* strain, which, after thorough washing, is plated as a mat on a synthetic agar medium containing a non-fermentable carbon source (e.g. ethanol) and no carnitine. On such plates, growing colonies of microorganisms that biosynthesise carnitine produce a zone due to the complementation of the Δ*cit2* mutant by carnitine. In this way, various microorganisms that produce carnitine were identified, including the yeasts *Yarrowia lipolytica*, *Rhodotorula graminis* and *Candida curvata* (Figure 3B). Endogenous carnitine biosynthesis by these strains was verified through intracellular carnitine/acetylcarnitine measurements using a novel HPLC-electrospray mass spectrometry (ESMS) method that is described above.

By adapting this large-scale screen, *S. cerevisiae* strains encoding functional BBH genomic fragments could be observed. As in the previous case, washed  $\Delta cit2$  cells were plated on a non-fermentable carbon source but in this instance,  $\gamma$ -butyrobetaine was added to the media to provide the necessary intermediate. Transformed wild type

strains forming zones would indicate functionally expressed BBH due to the formation of carnitine from γ-butyrobetaine and its subsequent secretion into the growth medium. Expressing the 2013 bp genomic fragment from the hypothetical protein NCU06891.1, resulted in the formation of a zone (Figure 3A). It was thus concluded that the gene encoded a BBH, and therefore the gene was named *cbs-2* for "carnitine biosynthesis gene no. 2".

#### Complementation of the carnitine-dependent \( \Delta cit2 \) strain by BBH

5

10

15

20

25

30

The pHVXII-cbs-2 construct encoding the 671 aa BBH homologue was transformed into FY23Acit2 in order to see if the transformed strains were able to grow on medium containing the precursor y-butyrobetaine. The transformants were streaked on synthetic glycerol media with and without y-butyrobetaine. Strain FY23\(\Delta\)cit2 transformed with the cbs-2 construct, grew in the presence of y-butyrobetaine, whereas the FY23\(\textit{Little of the media tested}\) the FY23\(\textit{Little of the media tested}\) (Figure 4A). Wild type strain FY23 grew normal on both glycerol media tested. The growth effect on glycerol agar plates were also clearly demonstrated on glycerol liquid media where the FY23\(\textit{\alpha}cit2\) strain transformed with the cbs-2 construct grew almost like the wild type strain and the FY23\(Delta\)cit2 strain transformed with the vector alone did not grow (Figure 4B). Interestingly, the FY23 wild type transformed with cbs-2 grew slightly better than the FY23 wild type transformed with vector alone indicating that production of carnitine can be advantageous for the cell or that γ-butyrobetaine may be slightly toxic to the cell. These data suggest that the γ-butyrobetaine in the growth medium is taken up by the  $\Delta cit 2$  strain and converted to carnitine endogenously, which allows the carnitine shuttle to function and therefore promote the production of energy and subsequent growth. To confirm this conclusion, carnitine and acetylcarnitine measurements were done using ESMS. Intra-cellular carnitine measurements were made after wild type strains were grown on synthetic medium containing glycerol and synthetic medium containing glycerol with y-butyrobetaine. No carnitine could be measured in FY23 wild type and FY23 transformed with cbs-2 in synthetic glycerol medium. When y-butyrobetaine was added, the FY23 transformed with cbs-2 gene showed high amounts of carnitine and acetylcarnitine, indicating that carnitine was produced and the carnitine shuttle was active (Figure 5). No carnitine could be measured in the FY23 wild type strain transformed with the vector alone, in the presence of y-butyrobetaine. A total of 897 ng/gWW of intracellular carnitine and 1151

ng/gWW intracellular acetylcarnitine was measured. S. cerevisiae only has carnitine acetyltransferase activity, so acetylcarnitine is the only carnitine ester that can be formed (Swiegers et al., 2001). Carnitine production of cbs-2 transformed strains could also be confirmed in glucose containing medium.

5

10

15

#### Discussion

In this study, a *S. cerevisiae* strain was genetically engineered that could biosynthesise carnitine from  $\gamma$ -butyrobetaine. A *N. crassa* genomic fragment was cloned that expressed a functional BBH, which could biosynthesise carnitine from  $\gamma$ -butyrobetaine. In addition, the BBH could suppress the growth defect of the carnitine-dependent  $\Delta cit2$  strain when cells were grown on glycerol synthetic medium containing  $\gamma$ -butyrobetaine.

The use of the large-scale selection screen could be useful to isolate carnitine overproducing mutants through monitoring the zone sizes. It can also be used to identify the novel carnitine biosynthesis genes from a variety of organisms. The use of carnitine producing strains of *S. cerevisiae* will increase the nutritional value of foods such as bread and beverages such as beer and wine. In addition, carnitine has recently been shown to protect *S. cerevisiae* from stress conditions (Lee et al., 2002). This would be an additional advantage to *S. cerevisiae* strains.

20

25

It will be apparent to those skilled in the art that other non-carnitine synthesising microorganisms, when transformed with this gene, would be able to synthesise carnitine from butyrobetaine, unless unable to transport this compound into the cell. This invention is therefore not intended to be limited to strains of *Saccharomyces cerevisiae*. Various alterations, modifications and other changes may also be made to the invention without departing from the spirit and scope of the present invention. It is therefore intended that the claims cover or encompass all such modifications, alterations and/or changes.

#### References

Becker, D. M. and Guarente, L. (1991). In *Guide to Yeast Genetics and Molecular Biology* (Guthrie, C., and Fink, G. R., eds), pp. 182-187, Academic Press, San Diego, CA. Bieber, L. L. (1988). Carnitine. *Annu Rev Biochem* 57, 261-83.

Bonnefont, J. P., Demaugre, F., Prip-Buus, C., Saudubray, J. M., Brivet, M., Abadi, N. and Thuillier, L. (1999). Carnitine palmitoyltransferase deficiencies. *Mol Genet Metab* 68, 424-40.

- Borum, P. R. and Broquist, H. P. (1977). Purification of S-adenosylmethionine: ε-N-L-lysine methyltransferase. The first enzyme in carnitine biosynthesis. *J Biol Chem* **252**, 5651-5.
- Bremer, J. (1983). Carnitine-metabolism and functions. Physiol Rev 63, 1420-80.
- Carter, A. L., Abney, T. O. and Lapp, D. F. (1995). Biosynthesis and metabolism of carnitine. *J Child Neurol* **10**, 3-7.
- Carter, H. E., Bhattacharyya, P. K., Weidman, K. R. and Fraenkel, G. (1952). Chemical studies on vitamin B<sub>T</sub> isolation and characterization as carnitine. *Arch Biochem Biophys* **38**, 405-16.
- Englard, S. (1979). Hydroxylation of γ-butyrobetaine to carnitine in human and monkey tissues. FEBS Lett 102, 297-300.
- Galland, S., Le Borgne, F., Bouchard, F., Georges, B., Clouet, P., Grand-Jean, F. and Demarquoy, J. (1999). Molecular cloning and characterization of the cDNA encoding the rat liver γ-butyrobetaine hydroxylase. *Biochim Biophys Acta* **1441**, 85-92.
- Hulse, J. D. and Henderson, L. M. (1980). Carnitine biosynthesis. Purification of 4-N-trimethylaminobutyraldehyde dehydrogenase from beef liver. *J Biol Chem* **255**, 1146-51.
- Kaufman, R. A. and Broquist, H. P. (1977). Biosynthesis of carnitine in *Neurospora crassa*. *J Biol Chem* **252**, 7437-9.
- Kelly, G. S. (1998). L-Carnitine: therapeutic applications of a conditionally-essential amino acid. Altern Med Rev 3, 345-60.
- Kunau, W. H., Buhne, S., De La Garza, M., Kionka, C., Mateblowski, M., Schultz-Borchard, U. and Thieringer, R. (1988). Comparative enzymology of β-oxidation. *Biochem Soc Trans* 16, 418-20.
- Lahjouji, K., Mitchell, G. A. and Qureshi, I. A. (2001). Carnitine transport by organic cation transporters and systemic carnitine deficiency. *Mol Genet Metab* **73**, 287-97.
- Lee, J., Lee, B., Shin, D., Kwak, S. S., Bahk, J. D., Lim, C. O. and Yun, D. J. (2002). Carnitine uptake by AGP2 in yeast Saccharomyces cerevisiae is dependent on Hog1 MAP kinase pathway. Mol Cells 13, 407-12.
- Lindstedt, G. and Lindstedt, S. (1970). Cofactor requirements of γ-butyrobetaine hydroxylase from rat liver. J Biol Chem 245, 4178-86.
- Pons, R. and De Vivo, D. C. (1995). Primary and secondary carnitine deficiency syndromes. J Child Neurol 10, 8-24.
- Rebouche, C. J. and Engel, A. G. (1980). Tissue distribution of carnitine biosynthetic enzymes in man. *Biochim Biophys Acta* **630**, 22-9.

Swiegers, J. H., Dippenaar, N., Pretorius, I. S. and Bauer, F. F. (2001). Carnitine-dependent metabolic activities in *Saccharomyces cerevisiae*: three carnitine acetyltransferases are essential in a carnitine-dependent strain. *Yeast* 18, 585-95.

- Swiegers, J. H., Vaz, F. M., Pretorius, I. S., Wanders, R. J. and Bauer, F. F. (2002). Carnitine biosynthesis in *Neurospora crassa*: identification of a cDNA coding for s-N-trimethyllysine hydroxylase and its functional expression in *Saccharomyces cerevisiae*. *FEMS Microbiol Lett* 210, 19-23.
- Van Roermund, C. W., Elgersma, Y., Singh, N., Wanders, R. J. and Tabak, H. F. (1995). The membrane of peroxisomes in *Saccharomyces cerevisiae* is impermeable to NAD(H) and acetyl-CoA under *in vivo* conditions. *EMBO J* 14, 3480-6.
- Van Roermund, C. W., Hettema, E. H., Van Den Berg, M., Tabak, H. F. and Wanders, R. J. (1999). Molecular characterization of carnitine-dependent transport of acetyl-CoA from peroxisomes to mitochondria in *Saccharomyces cerevisiae* and identification of a plasma membrane carnitine transporter, Agp2p. *EMBO J* 18, 5843-52.
- Vaz, F. M., Fouchier, S. W., Ofman, R., Sommer, M. and Wanders, R. J. (2000). Molecular and biochemical characterization of rat γ-trimethylaminobutyraldehyde dehydrogenase and evidence for the involvement of human aldehyde dehydrogenase 9 in carnitine biosynthesis. *J Biol Chem* **275**, 7390-4.
- Vaz, F. M., Ofman, R., Westinga, K., Back, J. W. and Wanders, R. J. (2001). Molecular and biochemical characterization of rat ε-N-trimethyllysine hydroxylase, the first enzyme of carnitine biosynthesis. *J Biol Chem* **276**, 33512-7.
- Vaz, F. M., Van Gool, S., Ofman, R., Ijlst, L. and Wanders, R. J. (1998). Carnitine biosynthesis: identification of the cDNA encoding human γ-butyrobetaine hydroxylase. *Biochem Biophys Res Commun* **250**, 506-10.
- Vaz, F. M., Van Gool, S., Ofman, R., Ijlst, L. and Wanders, R. J. (1999). Carnitine biosynthesis. Purification of γ-butyrobetaine hydroxylase from rat liver. *Adv Exp Med Biol* **466**, 117-24.
- Vaz, F. M. and Wanders, R. J. (2002). Carnitine biosynthesis in mammals. *Biochem J* 361, 417-29.
- Volschenk, H., Viljoen, M., Grobier, J., Petzold, B., Bauer, F., Subden, R. E., Young, R. A., Lonvaud, A., Denayrolles, M. and van Vuuren, H. J. (1997). Engineering pathways for malate degradation in Saccharomyces cerevisiae. Nat Biotechnol 15, 253-7.
- Winston, F., Dollard, C. and Ricupero-Hovasse, S. L. (1995). Construction of a set of convenient Saccharomyces cerevisiae strains that are isogenic to S288C. Yeast 11, 53-5.

## CLAIMS:

1. A method of producing a micro-organism that can biosynthesise carnitine, the method including the step of:

transforming a non-carnitine synthesising micro-organism with a nucleotide sequence encoding  $\gamma$ -butyrobetaine hydroxylase (BBH).

- 2. A method according to claim 1, wherein the micro-organism is a yeast strain.
- 3. A method according to claim 2, wherein the yeast strain is Saccharomyces cerevisiae.
- 4. A method according to any one of claims 1 to 3, wherein the transformed strain is cultured in the presence of  $\gamma$ -butyrobetaine in order for the strain to produce carnitine.
- 5. A method according to any one of claims 1 to 4, wherein the  $\gamma$ -butyrobetaine hydroxylase is a *Neurospora crassa*  $\gamma$ -butyrobetaine hydroxylase.
- 6. A method according to any one of claims 1 to 5, wherein the nucleotide sequence encoding  $\gamma$ -butyrobetaine hydroxylase (BBH) is a genomic fragment having the nucleotide sequence shown in Figure 7 (SEQ ID NO: 1).
- 7. A method according to any one of claims 3 to 6, wherein the Saccharomyces cerevisiae strain is a FY23 strain.
- 8. A method according to any one of claims 1 to 7, wherein the non-carnitine synthesising micro-organism is identified by:

applying the micro-organism to a synthetic culture medium containing a non-fermentable carbon source and no carnitine, and which is coated with a layer of Saccharomyces cerevisiae \( \Delta \) it2 strain;

culturing the micro-organism; and

detecting no formation of a zone produced in the culture medium in the region of the micro-organism if the micro-organism is not able to produce carnitine.

 A method according to any one of claims 1 to 8, wherein the success of producing a micro-organism that can biosynthesise carnitine is determined by

applying the transformed micro-organism to a synthetic culture medium containing a non-fermentable carbon source and no carnitine, and which is coated with a layer of Saccharomyces cerevisiae \( \Delta cit 2 \) strain;

culturing the micro-organism; and detecting the formation of a zone produced in the culture medium in the region of the micro-organism if the micro-organism is able to produce carnitine.

- 10. A strain of Saccharomyces cerevisiae that can produce carnitine in the presence of  $\gamma$ -butyrobetaine, which has been transformed with a nucleotide sequence encoding  $\gamma$ -butyrobetaine hydroxylase (BBH).
- 11. A Saccharomyces cerevisiae strain according to claim 10, wherein the γ-butyrobetaine hydroxylase is a Neurospora crassa γ-butyrobetaine hydroxylase.
- 12. A Saccharomyces cerevisiae strain according to either of claims 10 or 11, wherein the nucleotide sequence encoding  $\gamma$ -butyrobetaine hydroxylase (BBH) is a genomic fragment having the nucleotide sequence shown in Figure 7 (SEQ ID NO: 1).
- 13. A Saccharomyces cerevisiae strain according to any one of claims 10 to 12, wherein the Saccharomyces cerevisiae strain is a FY23 strain.
- 14. A method of identifying a carnitine-producing micro-organism, the method including the steps of:

applying a micro-organism to a synthetic culture medium which contains a non-fermentable carbon source and no carnitine, and which is coated with a layer of Saccharomyces cerevisiae \( \Delta cit2 \) strain;

culturing the micro-organism; and

detecting formation of a zone produced in the culture medium in the region of the micro-organism if the micro-organism is a carnitine-producing micro-organism.

15. A method according to claim 14, wherein the culture medium is illuminated or transilluminated in order to detect formation of the zone.

16. A method according to either of claims 14 or 15, wherein the culture medium is agar.

- 17. A method according to any one of claims 14 to 16, wherein the non-fermentable carbon source is ethanol.
- 18. A method according to claim 17, wherein the culture medium contains 2% ethanol synthetic agar medium.
- 19. A method according to any one of claims 14 to 18, wherein the micro-organism is cultured for a period of about 10 days at about 30 °C.
- 20. A method of producing carnitine, the method including the step of culturing in the presence of  $\gamma$ -butyrobetaine hydroxylase (BBH) a micro-organism that has been transformed with a nucleotide sequence encoding  $\gamma$ -butyrobetaine hydroxylase (BBH).
- 21. A method according to claim 20, wherein the micro-organism is included in the production of a beverage or food product.
- 22. A method of enhancing the nutritional value of a beverage or food product, the method including the step of incorporating a micro-organism that has been transformed with a nucleotide sequence encoding γ-butyrobetaine hydroxylase (BBH) in the process of manufacturing the food or beverage product.

Figure 1

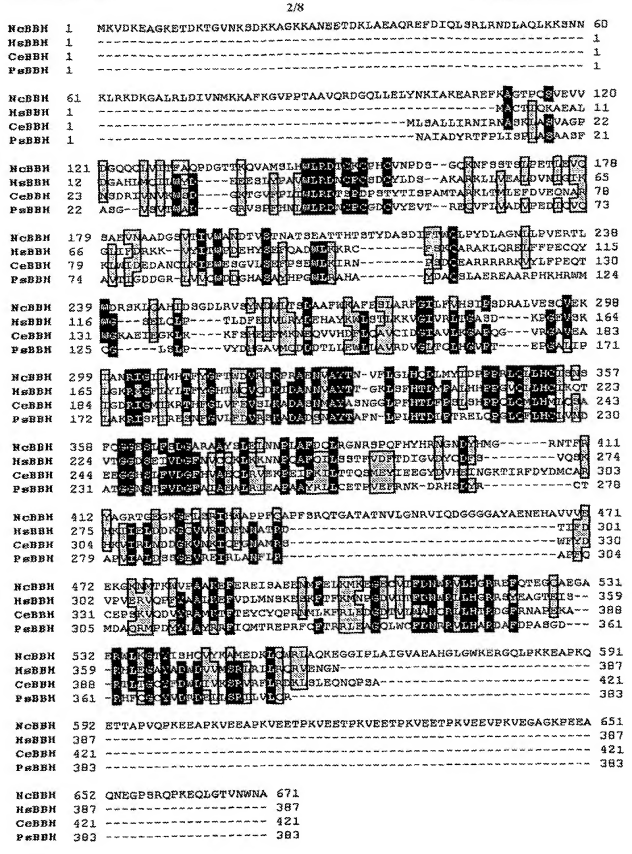
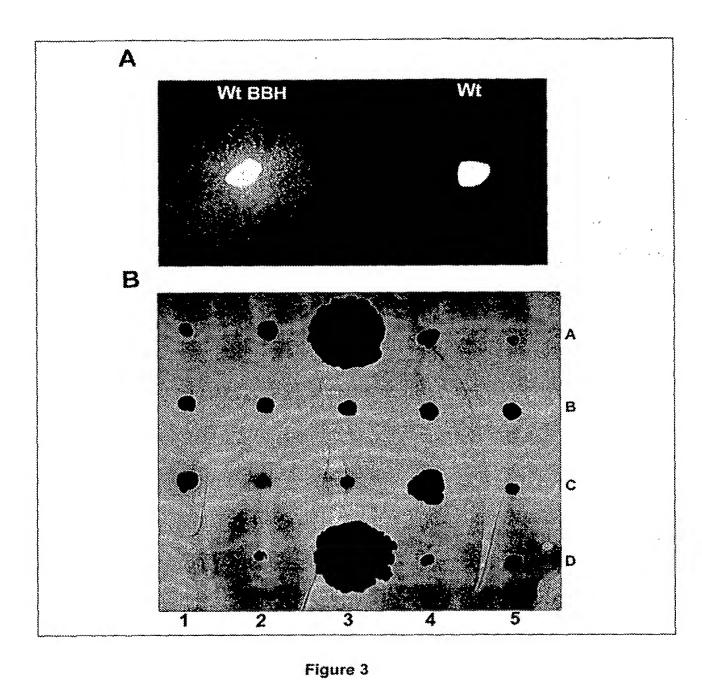


Figure 2



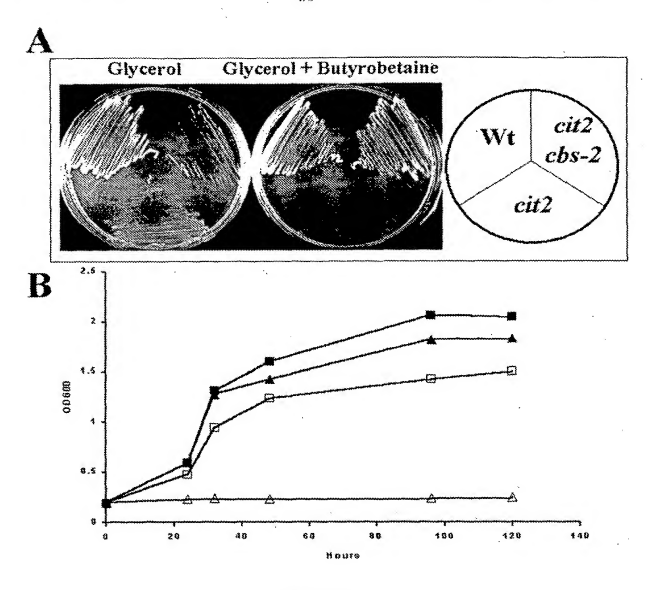


Figure 4

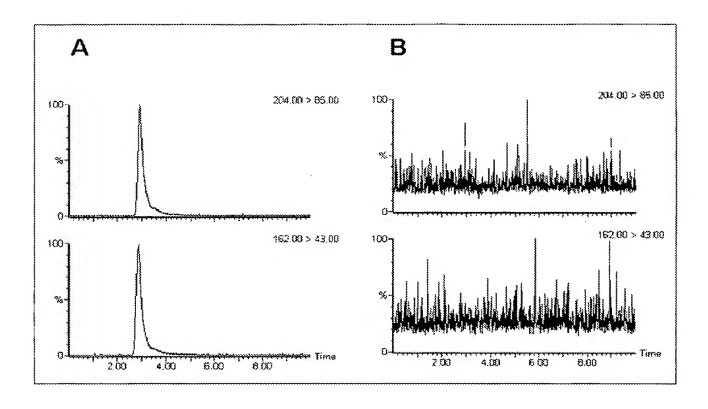


Figure 5

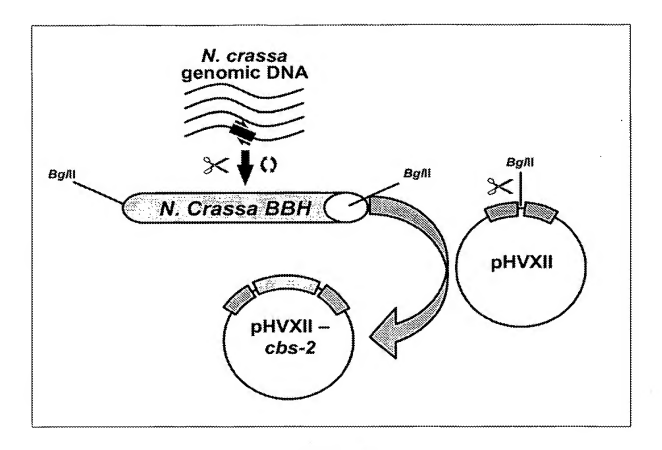


Figure 6

>NcBBH, 2016 bp

ATGAAAGTCGACAAGGAAGCCGGCAAGGAAACAGACAAGACCGGCGTCAA TAAATCGGACAAAAAGCCGGCAAGAAAGCCAACGAAGAAACGGACAAAT TGGCAGAAGCCCAGAGGGAATTCGATATACAACTCAGCCGGTTACGAAAC GACCTCGCCCAGCTAAAGAAATCCAACAACAAGTTGCGAAAAGACAAGGG GGCCTTACGGCTAGACATTGTCAATATGAAGAAAGCTTTCAAAGGGGTGC CTCCTACGCCAGCGGTGCAGAGGGACGGCCAATTGCTAGAATTATACAAC AAGATCGCCAAAGAAGCACGCGAATTCAAGGCCGCACTCCCCAGTCCGT CGAGGTGGTCGATGGTCAGCAGCAGCTCGTCATCACCTTCGCCCAGCCCG ACGGCACCACGAGCAAGTGGCCATGTCCCTCCACTGGCTGCGCGACACC TGCAAGTGCCCGCACTGCGTGAACCCCGACTCGGGCCAAAAGAACTTCTC CAGCACCTCTCTGCCGAGACTCTCGAGGTCCAAAGCGCCGAGGTCAACG CCGCCGACGCTCCGTCACCATCGTCTGGGCCAACGACACCGTCAGCACC AACGCCACGTCCGAGGCCACCACGCACACCTCGACCTACGACGCCTCCGA CATCTTTACCTGGCAACTTCCGTACGACCTCGCCGGCAACCTGCTCCCCG TCGAGCGCACGCTCTGGGACCGCTCCAAGCTGCAAGCCCACATCGACTCG GGCGACCTGCGCGTCTCGTACAACGACTGGCTGACCTCCGATGCCGCCTT TTGGAAAGCCTTTGAGTCTCTCGCCCGCTTCGGCATTCTCTTCGTGCACT CGATCCCGTCCGACCGAGCCCTCGTCGAATCCCAAGTCGAAAAGATCGCC AACCGCATCGGCATCCTCATGCACACCTTCTACGGCTTCACCTGGGACGT GCGCTCCAAGCCTCGCGCCGAGAACGTGGCCTACACCAACGTCTTTCTAG GCTTGCACCAAGACCTGATGTACATCGACCCGCCTCCCCGCCTGCAGCTC CTGCACTGCATCTCCAACTCCTTCCAGGGGGGGGAATCTCTCTTCAGCGA CGGAGCGCGCGCCTTACTCCCTGGAACTCAACAACCCACTAGCCTTTG ACCAGCTGCGCGCAACCGCTCGCCGCAGTTCCACTACCACCGCAACGGC AACGACTACCACATGGGCCGCAACACGTTCCGGTACGCCGGGCGGACGGG CGAAGGCAAGGGGTTTCTGAGCCGGATCCACTGGGCGCCGCCGTTCCAGG CGCCGTTTAGCCGGCAGACGGCGCCCACGGCGACGAACGTGCTCGGCAAC CGCGTCATTCAGGACGGCGGCGGCGGTGCTTATGCCGAGAACGAGCATGC GGTGGTGGTGGAGAAAAGGGCAAGAACATGACCAAGTGGGTGCCGGCGG CCAAGGAGTTTGAGCGCGAGATTAGCGCCGAGGAGAACATGTTTGAGCTC AAGATGAAGGAAGGAGTGTGTGATTTTCGATAACTGGCGAGTGTTGCA TGGGCGCAGGGAGTTCCAGACGGAAGGACAGGCGGAGGGCGCCGAGAGGT GGCTCAAGGGCACATATATTAGCCATCAGGTGTACAAGGCCATGGAGGAT AAGTTGCAGTGGAGGTTGGCGCAGAAGGAAGGGGGGATTCCTTTGGCTAT TGGCGTGGCGGAGGCGCATGGGTTGGGCTGGAAGGAGAGGGGGCAGTTGC CCAAGAAGGAGGCTCCTAAGCAGGAGACTACTGCCCCTGTTCAGCCCAAG GAGGAGCACCCAAGGTCGAGGAGGCACCCAAGGTCGAGGAGACTCCCAA GGTTGAGGAGACTCCCAAGGTTGAGGAGACTCCCAAGGTTGAGGAGACTC CCAAGGTCGAAGAGGTTCCCAAGGTCGAGGGGCCCGGGAAGCCCGAGGAG GCTCAGAATGAGGGTCCTTCGCGCCAGCCTAAGGAGCAATTGGGCACGGT GAACTGGAACGCATAA

Figure 7

NCBBH, 671 aa

MKVDKEAGKETDKTGVNKSDKKAGKKANEETDKLAEAQREFDIQLSRLRN

DLAQLKKSNNKLRKDKGALRLDIVNMKKAFKGVPPTAAVQRDGQLLELYN

KIAKEAREFKAGTPQSVEVVDGQQQLVITFAQPDGTTKQVAMSLHWLRDT

CKCPHCVNPDSGQKNFSSTSLPETLEVQSAEVNAADGSVTIVWANDTVST

NATSEATTHTSTYDASDIFTWQLPYDLAGNLLPVERTLWDRSKLQAHIDS

GDLRVSYNDWLTSDAAFWKAFESLARFGILFVHSIPSDRALVESQVEKIA

NRIGILMHTFYGFTWDVRSKPRAENVAYTNVFLGLHQDLMYIDPPRLQL

LHCISNSFQGGESLFSDGARAAYSLELNNPLAFDQLRGNRSPQFHYHRNG

NDYHMGRNTFRYAGRTGEGKGFLSRIHWAPPFQAPFSRQTGATATNVLGN

RVIQDGGGGAYAENEHAVVVEEKGKNMTKWVPAAKEFEREISAENMFEL

KMKEGECVIFDNWRVLHGRREFQTEGQAEGAERWLKGTYISHQVYKAMED

KLQWRLAQKEGGIPLAIGVAEAHGLGWKERGQLPKKEAPKQETTAPVQPK

EEAPKVEEAPKVEETPKVEETPKVEETPKVEEVPKVEGAGKPEE

AQNEGPSRQPKEQLGTVNWNA

Figure 8

60

#### PA137070-PCT as filed.ST25 SEQUENCE LISTING

<110> University of Stellenbosch <120> Method of producing a carnitine synthesising micro-organism <130> PA137070/PCT <150> ZA 2004/9060 <151> 2004-11-09 <160> 9 <170> PatentIn version 3.1 <210> 1 <211> 2016 <212> DNA <213> Neurospora crassa <400> 1 atgaaagtcg acaaggaagc cggcaaggaa acagacaaga ccggcgtcaa taaatcggac 120 aaaaaagccg gcaagaaagc caacgaagaa acggacaaat tggcagaagc ccagagggaa ttcgatatac aactcagccg grtacgaaac gacctcgccc agctaaagaa atccaacaac 180 240 aagttgcgaa aagacaaggg ggccttacgg ctagacattg tcaatatgaa gaaagctttc 300 aaaggggtgc ctcctacggc agcggtgcag agggacggcc aattgctaga attatacaac aagatcgcca aagaagcacg cgaattcaag gccggcactc cccagtccgt cgaggtggtc 360 garggreage ageageregt cateacerte geceagereg aeggeaceae caageaagtg 420 480 gccatgtccc tccactggct gcgcgacacc tgcaagtgcc cgcactgcgt gaaccccgac tegggeraaa agaacttete cageacetet etgeeegaga etetegaggt ecaaagegee 540 600 gaggicaacg cogeogacgg cicogicace atogicitggg coaacgacac ogicageacc 660 aacgccacgt cogaggccac cacgcacacc togacctacg acgcctccga catctttacc 720 togcaactic cgtacgacci cgccggcaac ctgctccccg tcgagcgcac gctctgggac cgctccaagc tgcaagccca catcgactcg ggcgacctgc gcgtctcgta caacgactgg 780 840 etgacetecg atgregeett ttggaaagee tttgagtete tegecegett eggeattete 900 ttegtgcact cgatcccgtc cgaccgagcc ctcgtcgaat cccaagtcga aaagatcgcc aaccgcatcg gcatcctcat gcacaccttc tacggcttca cctgggacgt gcgctccaag 960 cetegegeeg agaacgtgge etacaccaac gtetteetag gettgeacca agacetgatg 1020 1080

ggcgaatctc tcttcagcga cggagcgcgc gccgcttact ccctggaact caacaaccca 1140 ctagectttg accagetgeg eggeaacege tegeogragt tecaetacea eegeaacgge 1200 aacgactacc acatgggccg caacacgttc cggtacgccg ggcggacggg cgaaggcaag 1260 gggtttctga gccggatcca ctgggcgccg ccgttccagg cgccgtttag ccggcagacg 1320 ggcgccacgg cgacgaacgt gctcggcaac cgcgtcattc aggacggcgg cggcggtgct 1380

tacatogaco egecteceeg congrageto etgeactgea tetecaacte ettecaggge

#### PA137070-PCT as filed.5T25

tatgccgaga acgagcatgc ggtggtggtg gaggaaaagg gcaagaacat gaccaagtgg 1440 gtgccggcgg ccaaggagtt tgagcgcgag attagcgccg aggagaacat gtttgagctc 1500 aagatgaagg aaggagagtg tgtgattttc gataactggc gagtgttgca tgggcgcagg 1560 gagttccaga cggaaggaca ggcggagggc gccgagaggt ggctcaaggg cacatatatt 1620 agccatcagg tgtacaaggc catggaggat aagttgcagt ggaggttggc gcagaaggaa 1680 ggggggattc ctrtggctat tggcgtggcg gaggcgcatg ggttgggctg gaaggagagg 1740 gggcagttgc ccaagaagga ggctcctaag caggagacta ctgcccctgt tcagcccaag 1800 gaggaggcac ccaaggtcga ggaggcaccc aaggtcgagg agactcccaa ggttgaggag 1860 actcccaagg ttgaggagac tcccaaggtt gaggagactc ccaaggtcga agaggttccc 1920 aaggtcgagg gggccgggaa gcccgaggag gctcagaatg agggtccttc gcgccagcct 1980 aaggagcaat toggcacggt gaactggaac gcataa 2016

<210> 2

<211> 671

<212> PRT

<213> Neurospora crassa

<400> 2 Met Lys Val Asp Lys Glu Ala Gly Lys Glu Thr Asp Lys Thr Gly Val Ash Lys Ser Asp Lys Lys Ala Gly Lys Lys Ala Ash Glu Glu Thr Asp Lys Leu Ala Glu Ala Gln Arg Glu Phe Asp Ile Gln Leu Ser Arg Leu Arg Asn Asp Leu Ala Gln Leu Lys Lys Ser Asn Asn Lys Leu Arg Lys 50 60 ASP LYS GTY ATA Leu Arg Leu ASP ITE VAT ASH MET LYS LYS ATA Phe 65 70 80 Lys Gly Val Pro Pro Thr Ala Ala Val Gln Arg Asp Gly Gln Leu Leu
85 Glu Leu Tyr Asn Lys Ile Ala Lys Glu Ala Arg Glu Phe Lys Ala Gly 100 110 Thr Pro Gln Ser Val Glu Val Val Asp Gly Gln Gln Leu Val Ile 115 120 125 Thr Phe Ala Gln Pro Asp Gly Thr Thr Lys Gln Val Ala Met Ser Leu 130 135 140 His Trp Leu Arg Asp Thr Cys Lys Cys Pro His Cys Val Asn Pro Asp 145 150 155 160

Ser Gly Gln Lys Asn Phe Ser Ser Thr Ser Leu Pro Glu Thr Leu Glu 165 170 175

Val Gin Ser Ala Giu Val Asn Ala Ala Asp Gly Ser Val Thr Ile Val 180 190

Trp Ala Asn Asp Thr Val Ser Thr Asn Ala Thr Ser Glu Ala Thr Thr

PA137070-PCT as filed.ST25

His Thr Ser Thr Tyr Asp Ala Ser Asp Ile Phe Thr Trp Gln Leu Pro 210 215 278 Tyr Asp Leu Ala Gly Asn Leu Leu Pro Val Glu Arg Thr Leu Trp Asp 225 230 235 arg Ser Lys Leu Gin Ala His Ile Asp ser Gly Asp Leu Arg Val Ser 245 250 255 Tyr Asn Asp Trp Leu Thr Ser Asp Ala Ala Phe Trp Lys Ala Phe Glu 260 270 Ser Leu Ala Arg Phe Gly Ile Leu Phe val His Ser Ile Pro Ser Asp 275 280 285 Arg Ala Leu Val Glu Ser Gln Val Glu Lys Ile Ala Asn Arg Ile Gly 290 300 Ile Leu Met His Thr Phe Tyr Gly Phe Thr Trp Asp Val Arg Ser Lys 305 310 320 Pro Arg Ala Glu Asn Val Ala Tyr Thr Asn Val Phe Leu Gly Leu His Gln Asp Leu Met Tyr Ile Asp Pro Pro Pro Arg Leu Gln Leu Leu His 340 345 350 Cys Ile Ser Asn Ser Phe Gln Gly Gly Glu Ser Leu Phe Ser Asp Gly 360 365 Ala Arg Ala Ala Tyr Ser Leu Glu Leu Asn Asn Pro Leu Ala Phe Asp 370 380 Gln Leu Arg Gly Asn Arg Ser Pro Gln Phe His Tyr His Arg Asn Gly 385 390 395 Asn Asp Tyr His Met Gly Arg Asn Thr Phe Arg Tyr Ala Gly Arg Thr 405 415 Gly Glu Gly Lys Gly Phe Leu Ser Arg Ile His Trp Ala Pro Pro Phe 420 430 Gln Ala Pro Phe Ser Arg Gln Thr Gly Ala Thr Ala Thr Asn Val Leu 435 440 Gly Asn Arg val ile Gln Asp Gly Gly Gly Gly Ala Tyr Ala Glu Asn 450 460 Glu His Ala Val Val Glu Glu Lys Gly Lys Asn Met Thr Lys Trp 465 470 475 Val Pro Ala Ala Lys Glu Phe Glu Arg Glu Ile Ser Ala Glu Glu Asn 485 490 495 Met Phe Glu Leu Lys Met Lys Glu Gly Glu Cys Val Ile Phe Asp Asn 500 510 Trp Arg Val Leu His Gly Arg Arg Glu Phe Gln Thr Glu Gly Gln Ala 515 525 Glu Gly Ala Glu Arg Trp Leu Lys Gly Thr Tyr Ile 5er His Gln Val

									m a 1	ימכנו	in ne	~~ ~	. #4°	hal	2575		
Tyr 545	Ly5	Ala	Met	Glu	ASP 550	Lys	Fen	Gln	Trp	13707 Arg 555	Leu	Ala	GÌn	Lys	G1u 560		
Gly	Gly	Ile	Pro	Leu 56S	Ala	Ile	Gly	Val	A]a 570	Glu	Ala	H15	GTy	Leu 575	Gly		
тгр	Lys	Glu	Arg 580	бТу	Gln	Leu	Pro	Lys 585	Lys	Glu	Ala	Pro	Lys 590	GÌn	Glu		
Thr	Thr	Ala 595	Pro	Val	Gln	pro	Lys 600	Glu	Glu	Ala	Pro	Lys 605	val	Glu	Glu		
sΓα	Pro 610	Lys	val	Glu	Glu	Thr 615	Pro	Ly5	Vał	Glu	Glu 620	Thr	Pro	Lys	val		
G1u 625	Glu	Thr	pro	Lys	Va 1 630	Glu	Glu	Thr	Pro	Lys 635	val	Glu	61u	val	Pro 640		
Lys	/a?	Glu	G]y	A1a 645	GTy	i.ys	919	Glu	G1u 650	Ala	Gln	Asn	Glu	G1y 655	Pro		
Ser	Arg	Gln	Pro 660	Lys	Glu	Gln	Leu	գ <b>շ</b> 665	The	va]	Asn	Trp	Asn 670	Ala			
<21	0>	3															
<21		36															
<21		DNA															
<21	3>	Neui	rospc	ra c	rass	a											
<40 gat	0> caga	3 itct	atga	aagt	tog a	caag	gaaç	اد دؤ	ıgcaa	ı						36	
<21	.O>	4															
<23	1>	37															
<23	2>	DNA															
<21	13>	Neu	rospo	ora (	crass	sa											
<40 gat	)0> :cag:	4 atct	tta:	tgcg	ttc (	cagt	tcac	cg ti	gccc	вa						37	
<2	10>	5															
<2	11>	37															
<2	12>	ONA															
<b>&lt;2</b>	13>	Neu	rosp	ora	cras	sa											
<4 ga	00> tcga	S atto	atg	ctaa	ıgat	caaa	ttta	tg c	agag	ga						37	
<2	10>	6															
<2	11>	37															
<2	12>	DNA	<b>A</b>														
<2	13>	Net	ırosp	ora	cras	sa											
<4 ga	00> tcct	6 cga(	; tta	itttg	gtac	tgag	jgaa <i>ê</i>	ict t	crcr	itc		Pa	ge 4			37	,

#### PA137070-PCT as filed.ST25

```
<210> 7
<211>
<212>
      PRT
<213> Neurospora crassa
<400> 7
Pro Lys Val Glu Glu
<210> 8
<211> 3786
<212>
      DNA
<213> Neurospora crassa
<400>
atggggttcc tcgctactct catcgaccgt ggcatttccc acatcgacgg agcagcggcc
                                                                       60
agacgaggac ccaagtatag cccagacagc tactgcggcc gccgcctcgc cgggctcgaa
                                                                      120
acagggccga ttcccaaccg aggtcccgag acttcagccg ggctttggat cctcgtctcc
                                                                      180
ttctctacta ccttcctcgc tgcgaggtta tacctcaaga tgtaccgact gaaggggtta
                                                                      240
tggtgggatg attactitict tgttcttgct tggctcactc acaccctctc cgccaccctc
                                                                      300
geceaagtet ceateteest citeggeete ggceaetace cetgegaeat ecceteeeg
                                                                      360
acaacctcca tcccgcgcct gactctcgta ggcgaccact tcggcgccat gttctccatg
                                                                      420
ttcgccgtcg cgctctccaa gacttcctgg gccgtcactt tacttcggct tgttcgccgg
                                                                      480
                                                                      540
gggtctagta gtacctcctc ctcctcctcc tccaccacgt cctcttcttc ttccccggga
                                                                      600
caacaccage gecaataege agtetggatt gtetggtteg teateateae catetgtete
atcaagggcg ctcagggggt cttggtttgg attcccaaat gcggatcacc gcaggttgcg
                                                                      660
                                                                      720
cctgcagcat tgggggattc gaaagatgat aatgatgtgt cacatcatga tgtgtgtgtg
aggatcgaac cgctgaacgg gttcgcgacg ttcgcgggga gtgtcagtgg gacgtatgct
                                                                      780
attitgctgg cggtggtccc gtggaagacg attiggggca cgaattiggg caagagggaa
                                                                      840
awagitggag iggcgacgac gatgagcgig ggggcggiga giggggiggc ggcfiftgig
                                                                      900
ctggcggtca agatgaggag gattacgagt gagaatttta cttacgactc aggcgccatt
                                                                      960
                                                                     1020
atagtgtggt ctaccgccga gacgagcacg accatcatgg ctgcttgcat ccccgtccac
egegeettet geegeeaact tegeaagaag etgetggege agaacegett geatageage
                                                                     1080
aagccacact cgacgccgcc grcgacgggc aagaatggrg tcgggggtag tacgacarta
                                                                     1140
accaccciga acagiggicas citticatico giococigata agotticoca acaiggicigo
                                                                     1200
acgggaagtg attgcttgtc gttgtcttca ggtggtggag ggtgtcatgc cccgactgat
                                                                     1260
gacagogcaa gogacaaggo cattitigoag gitogigaca tigagagitig ogatataaat
                                                                     1320
gggcgtgagg tagttacctc aaggaacagt ggaaggatct tgaggacgca ggaggtcaag
                                                                     1380
gtggaatacc atcatcatcc tgagattttg gatgggaggg atgtgttgga tgagagggcg
                                                                     1440
cagtcacaac aagtcttcgt taacgccaag ttgtgcgccg acgccaaact cccgcttgta
                                                                     1500
cccgcatga agettggagt ggagctcatc cttggagccg ctcccctcgc gcgcggccgc
                                                                     1560
ttcqccqtca cgccatcaac caaggcgate cgcactttgt cgtgctcacc agcagctcgc
                                                                     1620
geogacgatg ctacgacccc ttctcccgcc gccggtgaca gtgccgcacc caccgagccg
                                                                     1680
                                                                     1740
qcccaqccca aactgctttt ccgtaagatc tataagaagc aaaggacagc ggaccagaaa
```

Page 5

## PA137070-PCT as filed.ST25

gccgacaaga	azggcgacaa	tggatccgac	atgaaagtcg	acaaggaagc	cggcaaggaa	1800
acagacaaga	ccggcgtcaa	taaatcggac	aaaaagccg	gcaagaaagc	caacgaagaa	1860
acggacaaat	tggcagaagc	ccagagggaa	ttcgatatac	aactcagccg	gttacgaaac	1920
gacctcgccc	agctaaagaa	atccaacaac	aagttgcgaa	aagacaaggg	ggccttacgg	1980
ctagacattg	tcaatatgaa	gaaagctttc	aaaggggtgc	ctcctacggc	agcggtgcag	2040
agggacggcc	aattgctaga	attatacaac	aagatcgcca	aagaagcacg	cgaattcaag	2100
gccggcactc	cccagtccgt	cgaggtggtc	gatggtcagc	agcagetegt	catcaccttc	2160
gcccagcccg	acggcaccac	caagcaagtg	gccatgtccc	tecactgget	gcgcgacacc	2220
tgcaagtgcc	cgcactgcgt	gaaccccgac	tcgggccaaa	agaacttctc	cagcacctct	2280
ctgcccgaga	ctctcgaggt	ccaaagcgcc	gaggtcaacg	ccgccgacgg	ctccgtcacc	2340
atcgtctggg	ccaacgacac	cgtcagcacc	aacgccacgt	ccgaggccac	cacgcacacc	2400
tcgacctacg	acgcctccga	catctttacc	tggcaacttc	cgtacgacct	caccaacage	2460
ctgctccccg	tcgagcgcac	gctctgggac	cgctccaagc	tgcaagccca	catcgactcg	2520
ggcgacctgc	gcgtctcgta	caacgactgg	ctgacctccg	atgccgcctt	ttggaaagcc	2580
tttgagtctc	tcgcccgctt	cggcattctc	ttcgtgcact	cgatcccgtc	egacegagee	2640
ctcgtcgaat	cccaagtcga	aaagategee	aaccgcatcg	gcatcctcat	gcacaccttc	2700
tacggcttca	cctgggacgt	gcgctccaag	cctcgcgccg	agaacgtggc	ctacaccaac	2760
gtctttctag	gcttgcacca	agacctgatg	tacatcgacc	cgcctccccg	cctgcagctc	2820
ctgcactgca	tctccaactc	crtccagggc	ggcgaatctc	tcttcagcga	cggagcgcgc	2880
gccgcttact	ccctggaact	caacaaccca	ctagcctttg	accagetgeg	cggcaaccgc	2940
tcgccgcagt	tccactacca	ccgcaacggc	aacgactacc	acatgggccg	caacacgttc	3000
cggtacgccg	ggcggacggg	rgaaggcaag	gggtttctga	gccggatcca	ctgggcgccg	3060
ccgttccagg	g cgccgtttag	ccggcagacg	ggcgccacgg	cgacgaacgt	gctcggcaac	3120
cgcgtcatto	aggacggcgg	cggcggtgct	tatgccgaga	acgagcatgo	ggtggtggtg	3180
gaggaaaagg	g gcaagaacat	t gaccaagtgg	grąccggcgg	ccaaggagtt	tgagcgcgag	3240
attagcgccg	g aggagaacat	t grtrgagete	aagatgaagg	aaggagagt	g tgtgattttc	3300
gataactgg	c gagtgttgc	a tgggcgcagg	gagttccaga	cggaaggac	a ggcggagggc	3360
gccgagagg	t ggctcaagg	g cacatatati	agccatcagg	tgtacaagg	c catggaggat	3420
aagttgcag	t ggaggttgg	c gcagaagga:	a ggggggatto	cttrggcta	t tggcgtggcg	3480
gaggcgcat	g ggttgggct	g gaaggagag	g gggcagttgc	ccaagaagg	a ggctcctaag	3540
					a ggaggcaccc	3600
aaggtcgag	g agactccca	a ggttgagga	g actoccaagg	g trgaggaga	c tcccaaggtt	3660
gaggagact	c ccaaggtcg	a agaggttcc	c aaggtcgagg	g gggccggga	a gcccgaggag	3720
gctcagaat	g agggtcctt	c gcgccagcc	t aaggagcaa	t tgggcacgg	t gaactggaac	3780
gcataa						3786

<210> 9

<211> 1261

<212> PRT

<213> Neurospora crassa

<400> 9

Met Gly Phe Leu Ala Thr Leu Ile Asp Arg Gly Ile Ser His Ile Asp 15

Page 6

PA137070-PCT as filed.ST25

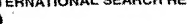
Gly Ala Ala Arg Arg Gly Pro Lys Tyr Ser Pro Asp Ser Tyr Cys 25 30 Gly Arg Arg Leu Ala Gly Leu Glu Thr Gly Pro Ile Pro Asn Arg Gly
40
45 Pro Glu Thr Ser Ala Gly Leu Trp Ile Leu Val Ser Phe Ser Thr Thr Phe Leu Ala Ala Arg Leu Tyr Leu Lys Met Tyr Arg Leu Lys Gly Leu 65 70 75Trp Trp Asp Asp Tyr Phe Leu Val Leu Ala Trp Leu Thr His Thr Leu 85 90 95 Ser Ala Thr Leu Ala Gln Val Ser Ile Ser Leu Phe Gly Leu Gly His 100 110 Tyr Pro Cys Asp Ile Pro Ser Pro Thr Thr Ser Ile Pro Arg Leu Thr Leu Val Gly Asp His Phe Gly Ala Met Phe Ser Met Phe Ala Val Ala 130 135 140 teu Ser tys Thr Ser Trp Ala Val Thr teu teu Arg Leu Val Arg Arg 145 150 160 Gly Ser Ser Ser Thr Ser Ser Ser Ser Ser Ser Thr Thr Ser Ser Ser 170 175 Ser Ser Pro Gly Gln His Gln Arg Gln Tyr Ala Val Trp 1le Val Trp 180 190 Phe Val Ile Ile Thr Ile Cys Leu Ile Lys Gly Ala Gln Gly Val Leu Val Trp Ile Pro Lys Cys Gly Ser Pro Gln Val Ala Pro Ala Ala Leu 210 215 220 Gly Asp Ser Lys Asp Asp Asp Asp Val Ser His His Asp Val Cys Val 725 230 235 Arg Ile Glu Pro Leu Asn Gly Phe Ala Thr Phe Ala Gly Ser Val Ser 250 255 Gly Thr Tyr Ala Ile Leu Leu Ala Val Val Pro Trp Lys Thr Ile Trp 265 270 Gly Thr Asn Leu Gly Lys Arg Glu Lys Val Gly Val Ala Thr Thr Met 275 280 Ser val Gly Ala val Ser Gly val Ala Ala Phe val Leu Ala val Lys 290 300 met arg arg Ile Thr Ser Glu Asn Phe Thr Tyr Asp Ser Gly Ala Ile 305 310 310 The val Trp Ser Thr Ala Glu Thr Ser Thr Thr Ile Met Ala Ala Cys Ile pro Val His arg Ala Phe Cys Arg Gin Leu Arg Lys Leu Leu 340

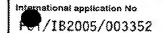
PA137070-PCT as filed.ST25
Ala Gln Asn Arg Leu His Ser Ser Lys Pro His Ser Thr Pro Pro Ser
355 360 365 Thr Gly Lys Asn Gly Val Gly Gly Ser Thr Thr Leu Thr Thr Leu Asn 370 Ser Gly Asn Phe His Ser Val Pro Asp Lys Leu Ser Gln His Gly Arg 385 390 395 Thr Gly Ser Asp Cys Leu Ser Leu Ser Ser Gly Gly Gly Gly Cys His
405
410 Ala Pro Thr Asp Asp Ser Ala Ser Asp Lys Ala Ile Leu Gin Val Arg ASP Ile Glu Ser Cys ASP Ile ASN Gly Arg Glu Val Val Thr Ser Arg 440 445 ASD Ser Gly Arg Ile Leu Arg Thr Glo Glu Val Lys Val Glu Tyr His His His Pro Glu Ile Leu Asp Gly Arg Asp Val Leu Asp Glu Arg Ala 465 470 470 Gin Ser Gin Gin Val Phe Val Asn Ala Lys Leu Cys Ala Asp Ala Lys
485
490 Leu Pro Leu Val Pro Arg Met Lys Leu Gly Val Glu Leu Ile Leu Gly 505 510 Ala Ala Pro Leu Ala Arg Gly Arg Phe Ala Val Thr Pro Ser Thr Lys Ala Ile Arg Thr Leu Ser Cys Ser Pro Ala Ala Arg Ala Asp Asp Ala 530 540 Thr Thr Pro Ser Pro Ala Ala Gly Asp Ser Ala Ala Pro Thr Glu Pro 545 550 555 Ala Gin Pro Lys Leu Leu Phe Arg Lys Ile Tyr Lys Lys Gin Arg Thr Ala Asp Gln Lys Ala Asp Lys Lys Gly Asp Asn Gly Ser Asp Met Lys 580 585 Val Asp Lys Glu Ala Gly Lys Glu Thr Asp Lys Thr Gly Val Asn Lys 595 600 605 Ser Asp Lys Lys Ala Gly Lys Lys Ala Ash Glu Glu Thr Asp Lys Leu 610 620 Ala Glu Ala Gln Arg Glu Phe Asp Ile Gln Leu Ser Arg Leu Arg Asn 625 630 635 ASP Leu Ala Gin Leu Lys Lys Ser Ash Ash Lys Leu Arg Lys Asp Lys Gly Ala Leu Arg Leu Asp Ile Val Ash Met Lys Lys Ala Phe Lys Gly
660 670 Val Pro Pro Thr Ala Ala Val Gln Arg Asp Gly Gln Leu Leu Glu Leu 675 680 685 Tyr Ash Lys Ile Ala Lys Glu Ala Arg Glu Phe Lys Ala Gly Thr Pro Page 8

PA137070-PCT as filed.ST25

Gin Ser Val Glu Val Val Asp Gly Gln Gln Gln Leu Val Ile Thr Phe Ala Gln Pro Asp Gly Thr Thr Lys Gln Val Ala Met Ser Leu His Trp 725 730 735 Leu Arg Asp Thr Cys Lys Cys Pro His Cys Val Asn Pro Asp Ser Gly 740 750 Gln Lys Asn Phe Ser Ser Thr Ser Leu Pro Glu Thr Leu Glu Val Gln
765 760 765 ser Ala Glu Val Asn Ala Ala Asp Gly Ser Val Thr Ile Val Trp Ala 770 780 Ash Asp Thr Val Ser Thr Ash Ala Thr Ser Glu Ala Thr Thr His Thr 785 790 795 800 Ser Thr Tyr Asp Ala Ser Asp Ile Phe Thr Trp Gln Leu Pro Tyr Asp 805 810 815 Leu Ala Gly Asn Leu Leu Pro Val Glu Arg Thr Leu Trp Asp Arg Ser 820 830 Lys Leu Gln Ala His Ile Asp Ser Gly Asp Leu Arg Val Ser Tyr Asn 840 845 ASP Trp Leu Thr Ser ASP Ala Ala Phe Trp Lys Ala Phe Glu Ser Leu 850 850 Ala Arg Phe Gly Ile Leu Phe Val His Ser Ile Pro Ser Asp Arg Ala 865 870 876 Leu Val Glu Ser Gln Val Glu Lys Ile Ala Asn Arg Ile Gly Ile Leu 885 890 Met His Thr Phe Tyr Gly Phe Thr Trp Asp Val Arg Ser Lys Pro Arg 900 910 Ala Glu Asn Val Ala Tyr Thr Asn Val Phe Leu Gly Leu His Gln Asp Leu Mer Tyr Ile Asp Pro Pro Pro Arg Leu Gln Leu Leu His Cys Ile 930 940 ser Asn Ser Phe Gln Gly Gly Glu Ser Leu Phe Ser Asp Gly Ala Arg 945 950 955 Ala Ala Tyr Ser Leu Glu Leu Asn Asn Pro Leu Ala Phe Asp Gln Leu 965 970 Arg Gly Asn arg Ser Pro Gln Phe His Tyr His Arg Asn Gly Asn Asp 980 980 Tyr His Met Gly Arg Asm Thr Phe Arg Tyr Ala Gly Arg Thr Gly Glu 995 1000 1005 Gly Lys Gly Phe Leu Ser Arg lle His Trp Ala Pro Pro Phe Glm 1010 1020 Ala Pro Phe Ser Arg Gln Thr Gly Ala Thr Ala Thr Asn Val Leu 1025 1030

- PA137070-PCT as filed.ST25 Gly Asn Arg Val Ile Gln Asp Gly Gly Gly Ala Tyr Ala Glu 1040 1050
- Asn Glu His Ala Val Val Glu Glu Lys Gly Lys Asn Met Thr 1055 1060 1065
- Lys Trp Val Pro Ala Ala Lys Glu Phe Glu Arg Glu Ile Ser Ala 1070 1080
- Glu Glu Asn Met Phe Glu Leu Lys Met Lys Glu Gly Glu Cys Val 1095
- The Phe Asp Ash Trp Arg val Leu His Gly Arg Arg Glu Phe Gln 1100 1110
- Thr Glu Gly Gln Ala Glu Gly Ala Glu Arg Trp Leu Lys Gly Thr
- Tyr Ile Ser His Gln Val Tyr Lys Ala Met Glu Asp Lys Leu Gln 1130 1140
- Trp Arg Leu Ala Gln Lys Glu Gly Gly Ile pro Leu Ala Ile Gly 1145 1155
- Val Ala Glu Ala His Gly Leu Gly Trp Lys Glu Arg Gly Gln Leu 1160 1170
- Pro Lys Lys Glu Ala Pro Lys Gln Glu Thr Thr Ala Pro Val Gln 1175 1180 1185
- Pro Lys Glu Glu Ala Pro Lys Val Glu Glu Ala Pro Lys Val Glu 1190 1200
- Glu Thr Pro Lys Val Glu Glu Thr Pro Lys Val Glu Glu Thr Pro 1205 1216
- Lys Val Glu Glu Thr Pro Lys Val Glu Glu Val Pro Lys Val Glu 1220 1230
- Gly Ala Gly Lys Pro Glu Glu Ala Gln Asn Glu Gly Pro Ser Arg 1245
- Gin Pro Lys Glu Gln Leu Gly Thr Val Asn Trp Asn Ala 1250 1260





A. CLASSIFICATION OF SUBJECT MATTER C12N1/16 C07K14/37 C12N9/02 C1201/34 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) C12N CO7K C120 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the injurnational search (name of data base and, where practical, search terms used) EPO-Internal, Sequence Search, EMBASE C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X VAZ F M ET AL: "Carnitine biosynthesis: 1-4,10 identification of the cDNA encoding human gamma-butyrobetaine hydroxylase." BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS. 18 SEP 1998, vol. 250, no. 2, 18 September 1998 (1998-09-18), pages 506-510, XP002365661 ISSN: 0006-291X page 507, column 2, paragraph 2 - page 509, paragraph 2 5-7, 11-13 -/--X X Further documents are listed in the continuation of Box C. See patent family annex. Special categories of cited documents : "T" tater document published after the international filling date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the \*A\* document defining the general state of the lart which is not considered to be of particular relevance. invention \*E\* earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone fising date \*L\* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" (document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled in the art. "O" document referring to an oral disclosure, use, exhibition or other means \*P\* document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 21/03/2006 7 March 2006 Name and mailing address of the ISA/ Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 Deleu, L

International application No /182005/003352

***************************************	ILION). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	GALAGAN JAMES E ET AL: "The genome sequence of the filamentous fungus Neurospora crassa." NATURE. 24 APR 2003, vol. 422, no. 6934, 24 April 2003 (2003-04-24), pages 859-868, XP002365858 ISSN: 0028-0836 the whole document & DATABASE UniProt [Online] 1 March 2004 (2004-03-01), "Hypothetical protein." retrieved from EBI accession no. UNIPROT:Q7S3G2 Database accession no. Q7S3G2 the whole document	5-7, 11-13
¥	LI X ET AL: "Identification of a novel family of nonclassic yeast phosphatidylinositol transfer proteins whose function modulates phospholipase D activity and Sec14p-independent cell growth."  MOLECULAR BIOLOGY OF THE CELL. JUN 2000, vol. 11, no. 6, June 2000 (2000-06), pages 1989-2005, XP002365662 ISSN: 1059-1524 figure 5	8,9,14, 15
Y	SWIEGERS J H ET AL: "Carnitine-dependent metabolic activities in Saccharomyces cerevisiae: three carnitine acetyltransferases are essential in a carnitine-dependent strain."  YEAST (CHICHESTER, ENGLAND) MAY 2001, vol. 18, no. 7, May 2001 (2001-05), pages 585-595, XP002365859 ISSN: 0749-503X figure 1	8,9,14, 15
X	EP 0 410 430 A (LONZA AG) 30 January 1991 (1991-01-30) page 2, line 29 - line 33; claim 1 page 2, line 38	20
X	PRETORIUS I S ET AL: "Meeting the consumer challenge through genetically customized wine-yeast strains" TRENDS IN BIOTECHNOLOGY, ELSEVIER PUBLICATIONS, CAMBRIDGE, GB, vol. 20, no. 10, 1 October 2002 (2002-10-01), pages 426-432, XP004379928 ISSN: 0167-7799 page 431, column 1, paragraph 2	21,22

International application No /IB2005/003352

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT						
ategory*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.				
· , χ	WO 2005/083089 A (CJ CORPORATION; CHUNG, SUNG OH; LEE, BHEONG-UK; KANG, WHAN-KOO; JU, JA) 9 September 2005 (2005-09-09) the whole document	1,2,5				
		12				

Form PCT/ISA/210 (continuation of second sheet) (April 2005)

Information on patent family members

International application No FOI/IB2005/003352

Patent document cited in search report		Publication date		Patent family member(s)	Publication date	
EP 0410430	A	30-01-1991	AT	132535 T		
			AU	625525 B2	16-07-1992	
			AU	5983790 A	31-01-1991	
			BR	9003672 A	27-08-1991	
			DD	296702 A5	12-12-1991	
			DE	59010027 D1	15-02-1996	
			DK	410430 T3	29-01-1996	
			ES	2081878 T3	16-03-1996	
			FI	102083 B1	15-10-1998	
			IE	902689 A1	27-02-1991	
			IL	95196 A	26-05-1995	
			JP	2018232 C	19-02-1996	
			JP	3076591 A	02-04-1991	
			JP	7051071 B	05-06-1995	
			MX	170707 B	08-09-1993	
			NO	903338 A	29-01-1991	
WO 2005083089	A	09-09-2005	NONE		a report referent water states states states about 1900 about major jabor variar variar variar variar.	